Robust dicopper(I) μ-boryl complexes supported by a dinucleating naphthyridine-based ligand

Pablo Ríos, Matthew S. See, Rex C. Handford and T. Don Tilley

Copper boryl species have been widely invoked as reactive intermediates in Cu-catalysed C–H borylation reactions, but their isolation and study have been challenging. Use of the robust dicopper(I) boryl ligand, DPFN (2,7-bis(fluoro-di(2-pyridyl)methyl)-1,8-naphthyridine) allowed for the isolation of two very thermally stable dicopper(I) boryl complexes, [(DPFN)Cu₂(μ-Bpin)][NTf₂] (2) and [(DPFN)Cu₂(μ-Bcat)][NTf₂] (4) (pin = 2,3-dimethylbutane-2,3-diol; cat = benzene-1,2-diol). These complexes were prepared by cleavage of the corresponding diborane via reaction with the alkoxide [(DPFN)Cu₂(μ-O'Bu)][NTf₂] (3). Reactivity studies illustrated the exceptional stability of these boryl complexes (thermal stability in solution up to 100 °C) and their role in the activation of C(sp)−H bonds. X-ray diffraction and computational studies provide a detailed description of the bonding and electronic structures in these species, and suggest that the dicopper(I) character of the naphthyridine-based ligand is largely responsible for their remarkable stability.

Introduction

Copper-catalysed C–H borylation is a powerful tool in organic chemistry due to the stability and versatility of the resulting organoboron species as synthetic building blocks. This catalysis is thought to involve Cu(I) boryl (Cu–BR₂) complexes as key reactive intermediates. However, the chemistry of Cu(I) boryls has only recently been observed; while the first proposal of a Cu–BR₂ intermediate stems from a 2000 report by Miyaura and Hosomi, the first well-defined Cu(I) boryl species was described by Sadighi and coworkers in 2005. Since then, only a few have been isolated. The paucity of examples is likely due to their inherent instability, given that most of those reported decompose in solution at or below room-temperature to elemental Cu. Notably these cases involve monodentate N-heterocyclic carbene (NHC) or phosphine supporting ligands. In 2016, Sadighi reported the first dicopper boryl, [(SiPr)Cu₂(μ-Bcat)](BF₄) (Scheme 1, top left, SiPr = 1,3-bis(2,6-diisopropylphenyl)imidazol-2-ylidene), from reaction of [(SiPr)Cu₂(μ-OSiMe₃)](BF₄) with bis(catecholato)diboron (B₂cat₂) at −35 °C. The same dicopperation was obtained by Kleeberg in 2019 upon treatment of mononuclear (SiPr)CuOtBu with B₂cat₂ in THF. Interestingly, mononuclear (SiPr)CuBcat was observed when the reaction was carried out in toluene. By modifying the boryl and supporting ligands, the Kleeberg group obtained additional examples of neutral dicopper boryls (Scheme 1).

Scheme 1. Examples of dicopper(I) μ-boryl complexes

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solution at or below room-temperature and require storage in the solid-state under an inert atmosphere, complicating their characterisation and study as catalytic intermediates.

Here we report the synthesis and characterisation of two dicopper(II) boryl complexes supported by a 1,8-naphthyridine-based dinucleating ligand. These species are persistent in solution and tolerate heating up to at least 100 °C. In addition, their role in the borylation of C(sp)−H bonds and the origins of their stability are described.

**Results and discussion**

Previous results from this laboratory established 2,7-bis(fluoro-di(2-pyridyl)methyl)-1,8-naphthyridine (DPFN) as an effective platform for stabilisation of bimetallic units. Dicopper(II) [[DPFN]Cu$_2$] complexes, accessed via the convenient starting material [[DPFN]Cu$_2$(µ-NCMMe)][NTf$_2$] (1), exhibit remarkable bimetallic electrophilicity and have allowed for the study of several elusive reactive intermediates such as a dicopper(II) nitrenoid, pentanuclear metal hydrides, and an unprecedented dicopper(II) bridging triazolide (Scheme 2). Thus, [[DPFN]Cu$_2$] seemed a promising framework for the synthesis and stabilisation of dicopper boryl units.

Due to the electrophilic character of 1, introduction of a boryl fragment requires an anionic borylation reagent. Whereas few synthons for boryl anions are available, a Lewis base adduct of a tetraalkoxy diboron (e.g., alkoxide, fluoride or NHC) gives an isolable sp$^3$-sp$^3$ anionic diboron compound capable of transferring a boryl group to an electrophile under metal-free conditions. To pursue this possibility, one equivalent of K[B$_2$pin$_2$O$_2$Bu] was added to complex 1 in ortho-difluorobenzene (o-DFB) at 25 °C to give a color change from orange to dark green after 3 h. A $^1$H NMR analysis revealed formation of a single new symmetrical species (2) with a set of resonances corresponding to the DPFN ligand. This was corroborated by $^{19}$F($^1$H) NMR spectroscopy, which showed that the singlet at $-175.1$ ppm for 1 had been replaced by a singlet at $-177.8$ ppm. The $^1$H NMR spectrum of the reaction mixture revealed the presence of free MeCN and BuOBpin, as well as a singlet at 1.44 ppm associated with 2. This assignment is supported by the $^{11}$B NMR spectrum containing a single broad resonance for BuOBpin at 21.7 ppm, in agreement with reported values (Figures S13). Altogether, these data suggest the formation of boryl complex [[DPFN]Cu$_2$(µ-Bpin)][NTf$_2$] (2, Scheme 3).

Scale-up of the synthesis of 2, or use of THF as solvent led to formation of several side-products, as ascertained by $^1$H NMR analysis (Figure S14). It was hypothesized that one possible side product could result from the presence of free KO Bu, generated in solution by dissociation from B$_2$pin. To test this hypothesis, one equivalent of KO Bu was added to a solution of 1 in o-DFB at 25 °C, after which the reaction mixture became dark orange (Scheme 4).


Figure 1. Solid-state molecular structure of 3 (50% probability ellipsoids); H atoms are omitted for clarity. Only the major disorder component of the tert-butoxide fragment is shown.
a new set of resonances resulting from the DPFN ligand, along with a new singlet at 1.76 ppm, indicating the clean formation of ([DPFN]Cu$_3$(µ-O’Bu))[NTf$_2$] (3). The resonances for 3 correspond to those of one of the aforementioned side-products formed in the synthesis of 2, and confirms the release of KO’Bu in solution to some extent.

Complex 3 was isolated by precipitation from o-DFB or THF with pentane, but removal of volatile components from the resulting powder in vacuo resulted in conversion to an insoluble, intractable material. Nonetheless, dark brown X-ray quality crystals were grown by layer diffusion of O(SiMe$_3$)$_2$ into an o-DFB solution of 3 (generated in solution) after 3 days at 25 °C. The solid-state structure, as determined by single crystal diffraction analysis, confirms the proposed assignment (Figure 1). While the Cu–O bond distances are similar to those observed in previously reported dicopper complexes containing a 1,8-naphthyridine diphosphine ligand (ca. 1.9 Å), the Cu···Cu distance in 3 is much shorter (2.687(1) Å vs. 2.9-3.0 Å) and the Cu–O–Cu angle is considerably more acute (87.6(2)° vs 100-108°), indicating a closer metal-metal contact enforced by the dipyridyl (vs. phosphinyl) side-arms. However, the metrical parameters for 3 are in good agreement with previous results from this laboratory on the dicopper(II) DPFN complex with a bridging aryloxide ligand.

With a reliable route to complex 3, reactions with diboranes were explored. Addition of B$_2$pin$_2$ to complex 3 in THF (prepared in situ from 1) resulted in a dark green solution after stirring at 25 °C for 15 h. Clean formation of boryl species 2 and 1'BuOBpin (Scheme 5) was observed by $^1$H, $^{19}$F($^1$H) and $^{11}$B($^1$H) NMR spectroscopy, and no side-products were observed upon increasing the concentration of the reaction solution. Complex 2 was isolated as green crystals by layering pentane over a o-DFB/THF solution which gave analytically pure 2 in 75% yield. The identity of the complex (Figure 2, top) was confirmed by single-crystal X-ray diffraction. Similarly, B$_2$cat$_2$ reacted with complex 3 generated in situ to give the corresponding dicopper(I) boryl complex ([DPFN]Cu$_3$(µ-Bcat))[NTf$_2$] (4) after 2 h at 25 °C (Scheme 5), as judged by multinuclear NMR analysis on the resulting purple THF solution. The much shorter reaction time compared to that for the formation of 2 might be due to the higher Lewis acidity of B$_2$cat$_2$ in comparison to B$_2$pin$_2$. Similar work-up conditions afforded pure 4 as green X-ray quality crystals in 85% yield (Figure 2, bottom). Unlike most previously described mono- or dicopper boryl complexes, 2 and 4 were synthesised and isolated at 25 °C, indicating a considerably higher stability. This is also reflected in mass spectra (Figures S27-S28), which indicate persistence of the dinuclear structures under the ionisation conditions. The increased stability of these boryl complexes in solution raises interest in their bond metrics compared to those of reported, thermally unstable analogues. However, 2 and 4 possess geometrical parameters that are within the range of those reported for dicopper µ-boryl complexes. The Cu–B bond distances (2.06-2.09 Å) are slightly longer than those in cationic ([{SiPr}Cu$_2$(µ-Bcat)][NF$_4$] (ca. 2.05 Å) but are considerably shorter than those in the dinuclear complexes synthesised by Kleeberg and co-workers (> 2.17 Å). Similarly, 2 and 4 feature Cu–B–Cu bond angles (ca. 68°) that are more acute than that in Sadighi’s dinuclear complex (72.1(2)°) but more obtuse than in Kleeberg’s complexes (ca. 60°). Finally, the Cu···Cu distances (ca. 2.32 Å) are between those of the cationic (2.4083(9) Å) and neutral (ca. 2.22-2.27 Å) µ-boryl examples previously reported. Nevertheless, all such distances are significantly shorter than the sum of covalent radii for Cu (2.64 Å), implying that the short metal-metal distances are due to the bridging boryl ligands and/or cuprophilic interactions. Interestingly, the increased stability of 2 and 4 in solution does not seem to be attributable to peculiar binding metrics. The robust character of these boryl species is also manifested in...
their reactivity. Both 2 and 4 exhibited no reaction with a number of small molecules including internal alkynes, olefins, silanes, azides or organic molecules possessing weak C–H bonds (e.g. fluorene), over a range of reaction conditions (THF or o-DFB as solvents and temperatures up to 70 or 100 °C, respectively; Table S1(Figures S15-S16). In all cases, these bimetallic complexes coexist in solution with the added reagent even at reaction times of 24 h. In contrast to results reported by Kleeberg et al. for dicopper, 2 and 4 are inert towards 4-iodotoluene. Aldehydes (1,3,5-trioxane or mesitylaldehyde) did not react with 2 or 4, in stark contrast to results observed for (IPr)CuBpin (IPr = 1,3-bis(2,6-diisopropylphenyl)imidazol-2-ylidene), demonstrating how nuclearity can have a dramatic influence on the reactivity of boryl derivatives. Warming of 2 or 4 to 75 °C in the presence of either CO2 or O2 gave rise to a complex mixture of products, as shown in the NMR spectra of Figures S17-S18, whereas H2 was unreactive. In contrast, HClC(p-CF3-C6H4) cleanly reacted with both bridging boryl complexes to yield [(DPFN)Cu2(µ-C≡C(C6H4)CF3)][NTf2] (5), as summarised in Scheme 6 (Figures S19-S21). The increased Lewis basicity of boryl 2 compared to 4 is evident from the reaction conditions necessary to achieve high (>80%) conversions: whereas 2 reacted with 1.5 equivalents of alkene at 70 °C in 21 hours, boryl 4 required more forcing conditions (6 equivalents of alkylene and warming for 90 hours at 100 °C). Interestingly, bridging alkylene 5 further reacted with B2cat2 (but not with B2pin2) to regenerate Bcat complex 4 (Figure S21), and this reactivity trend is consistent with the observed reactions of 3 with diboranes (vide supra). The clean deprotonation of HC≡C(p-CF3-C6H4) to give 5 at high temperatures is surprising, given the potential side-reactions that are possible in the presence of free hydroborane, such as alkynecarbonypin → Cu. Bertrand and coworkers recently reported catalytic C(sp)–H dehydrogenative borylation using carbene-stabilised Cu complexes, presumably by way of a catalytically active α,π-dicopper acetylide intermediate A (Scheme 7A). Unlike species A, the triple bond of bridging alkynyl 5 is not engaged in π-bonding to copper, making it susceptible to hydroboration processes. Control experiments involved the independent synthesis of 5 by treating the previously reported bridging phenyl complex [(IPr)Cu2(µ-Ph)][NTf2]27 with HClC(p-CF3-C6H4). Addition of HBPin or HBcat to 5 did not result in reaction at 25 or 100 °C over 48 h (Figures S24-25). This divergence in reactivity from that observed by Bertrand et al. could be due to the different binding mode of the alkynyl fragment to the dicopper core (α,π vs α,σ), or to the rigidity imposed by the dinucleating DPFN ligand.

Computational methods were used to gain insights into the unusual robustness of bridging boryls 2 and 4. First, steric congestion around the boron atom was quantified by determining its percent buried volume (%Vbur), described as the volume of a sphere centered on boron that is occupied by the [(DPFN)Cu2]+ scaffold and the substituents on B (Figures S31-34). As expected, %Vbur values for 2 and 4 are remarkably large (ca. 80%), yet intermediate between values observed for unstable species in solution, namely mononuclear (IPr)CuBpin (%Vbur = 65.3%), and [(SiPr)Cu2(µ-Bcat)][BF4] (%Vbur = 83.4%).

<table>
<thead>
<tr>
<th>Parameter</th>
<th>[(DPFN)Cu2(µ-Ph)][NTf2]</th>
<th>[(IPr)Cu2(µ-Bpin)][NTf2]</th>
<th>[(IPr)Cu2(µ-Bcat)][BF4]</th>
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<tr>
<td>Nat. charge on Cu</td>
<td>0.32</td>
<td>0.31</td>
<td>0.37</td>
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<tr>
<td>Nat. charge on B</td>
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<td>0.59</td>
<td>0.50</td>
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<tr>
<td>Cu–B Bond order</td>
<td>0.65</td>
<td>0.67</td>
<td>0.58</td>
</tr>
<tr>
<td>Cu–Cu Bond order</td>
<td>0.73</td>
<td>0.72</td>
<td>0.32</td>
</tr>
</tbody>
</table>

Table 1. Natural charge and bond order values for selected computed dicopper(i) µ-boryl complexes. *Values obtained from reference 6.
In light of this, steric hindrance around boron is not likely to be a strong contributor to the stability. DFT analyses were performed on 2 and 4 to understand their electronic structures, using the PBE0-D3/6-31g(d,p)/SDD level of theory on the cationic fragments. The optimised structures were found to be local minima, with geometries in excellent agreement with metrics observed for the solid state structures (Table S2). Natural charges on Cu and B reflect minimal differences with those of related cationic systems, as displayed in Table 1. Likewise, Wiberg Bond Order analysis exhibited similar values for all Cu–B bonds, albeit the Cu−Cu interactions seem to be stronger in the cationic fragments of 2 and 4, as expected based on the observed, short Cu···Cu distances. In addition, no bond critical point between the Cu atoms was observed by QTAIM calculations (Figures S37-38). Finally, the electron distribution in the Cu–B bonds was investigated, since studies attribute the high nucleophilicity and reactivity exhibited by these complexes to the Cu-boryl σ-bonding electrons. Computational work by Carbó and Fernández in 2012, and Sheong and Lin in 2021, revealed that the contribution of B-based orbitals to the M−BR2 bond is remarkably high (ca. 70%, with a p/s ratio ≈ 0.9) in the case of mononuclear Cu-boryl species, whereas in other examples like Au- or Pd-boryl compounds, the contribution of B is around 56% and 53%, respectively. Therefore, the Cu−B bond tends to be substantially more polarised towards the B atom. Natural Localised Molecular Orbital (NLMO) analysis of 2 and 4 reveals σ-donation of the boryl ligand to the empty 4s orbitals on the Cu atoms (as previously observed for [(DPFN)Cu2]n complexes) with negligible back-donation from the metals to boron. This 3c-2e bonding scenario is illustrated in Figure 3. Surprisingly, the contributions of Cu- and B-based orbitals to the bonding are almost identical to those previously reported for reactive, mononuclear Cu-boryl complexes (ca. 30% Cu and 70% B), with a considerably smaller p/s ratio on B in the case of 4.

These data indicate that the extreme stability and chemical inertness of boryls 2 and 4 in polar solvent does not stem from peculiar geometrical or electronic factors, since these properties are similar to those of much less stable analogues. Rather, the rigid framework imposed by the dinucleating 1,8-naphthyridine-based ligand seems to be responsible for the observed stability, by precluding dynamic behaviour in solution that might produce an equilibrium concentration of more reactive, monomeric species. This hypothesis finds additional support from a recent study by Ito and coworkers, which concludes that a borylcopper(I) dimer is the dormant species in a catalytic asymmetric borylation process, whereas the mononuclear Cu-boryl complex is the active form of the catalyst (Scheme 8). 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Notes and references


17. The formation of an insoluble material after exposing complex 3 to vacuum prevented the synthesis of analytically pure solid samples of this compound.


24. Concomitant formation of HBpin is observed in the case of complex 4. In the case of 4, HBcat seems to decompose over time to catBOBcat under the reaction conditions. This has been confirmed by warming HBcat in o-DPB at 100 °C (see Figures S22 and S26 for more information).


30. The calculations were performed with the Gaussian 16 program: Gaussian 16, Revision A.03, Gaussian, Inc. Wallingford CT, 2016. The full Gaussian citation can be found in the ESI.


The p/s ratio also gives an idea of the nucleophilicity of a M-boryl bond. A boryl ligand with more p character on B will donate more electron density to the metal, making the M-B bond stronger. On the contrary, a boryl ligand with more s character on B will polarise the M-B bond towards B. For more information, see reference 34 or J. Zhu, Z. Lin and T. B. Marder, *Inorg. Chem.*, 2005, **44**, 9384-9390.